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         JUL 02
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         JUL 02
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         JUL 02
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                CAplus enhanced with French and German abstracts
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                 spectral property data
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                 BEILSTEIN updated with new compounds
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         NOV 15
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NEWS EXPRESS
             19 SEPTEMBER 2007: CURRENT WINDOWS VERSION IS V8.2,
              CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007.
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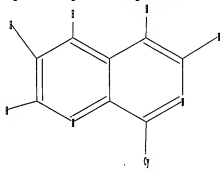
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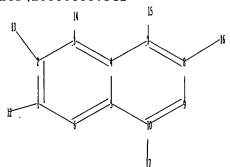
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chain nodes :

12 13 14 15 17

ring nodes :

1 2 3 4 5 6 7 8 9 10 16

chain bonds :

1-12 2-13 3-14 7-15 8-16 10-17

ring bonds :

1-2 1-6 2-3 3-4 4-5 4-7 5-6 5-10 7-8 8-9 9-10

exact/norm bonds :

8-16 10-17

exact bonds :

1-12 2-13 3-14 7-15

normalized bonds :

1-2 1-6 2-3 3-4 4-5 4-7 5-6 5-10 7-8 8-9 9-10

isolated ring systems :

containing 1 :

Match level:

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:Atom 17:Atom

L1 STRUCTURE UPLOADED

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L1 HAS NO ANSWERS

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100.0% PROCESSED 104 ITERATIONS

ITERATIONS 1 ANSWERS

SEARCH TIME: 00.00.01

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PROJECTED ITERATIONS:

1469 TO 2691

PROJECTED ANSWERS: 1 TO

L2 1 SEA SSS SAM L1

=> s l1 full

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FULL SCREEN SEARCH COMPLETED - 2232 TO ITERATE

100.0% PROCESSED 2232 ITERATIONS

64 ANSWERS

SEARCH TIME: 00.00.01

L3 64 SEA SSS FUL L1

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L41 L3

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ACCESSION NUMBER: 2004:534201 CAPLUS

DOCUMENT NUMBER: 141:71530

TITLE: Preparation of [1,7] naphthyridines as PDE4 inhibitors

INVENTOR(S): Denholm, Alastair; Keller, Thomas Hugo; Mccarthy,

Clive; Press, Neil John; Taylor, Roger John

PATENT ASSIGNEE(S): Novartis Ag, Switz.; Novartis Pharma GmbH

SOURCE: PCT Int. Appl., 38 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.					KIN	D	DATE			APPL:	I CAT	ION	иО.		D	ATE		
	2004	0550	1 2		71	-	2004	0701			002		262			0021	216	
WO	2004	0330.	13		ΑI		2004	0 / 0 1	1	WO 2	003-	CPI4.	203			0031	213	
	W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
		co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	GE,	
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		LU,	LV,	MA,	MD,	MK,	MN,	MX,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	
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		SI,	SK,	TR														
CA	2505	405			A1		2004	0701	1	CA 2	003-	2505	405		2	0031	215	
AU	2003	2938	86		A1		2004	0709		AU 2	003-	2938	86		2	0031	215	
EP	EP 1575950			A1	20050921		EP 2003-789283						2	0031	215			
	R:	AT,	BE,	CH,	DE,	DK.	ES,	FR,	GB,	GR,	IT.	LI,	LU.	NL.	SE.	MC.	PT.	

IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK BR 2003017330 Α 20051108 BR 2003-17330 20031215 CN 1726215 20060125 CN 2003-80106300 Α 20031215 JP 2006511539 Т JP 2004-560419 20060406 20031215 EP 1777226 EP 2007-100446 A1 20070425 20031215 AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LI, LU, MC, NL, PT, RO, SE, SI, SK, TR US 2006058338 **A**1 20060316 US 2005-538355 20050808 PRIORITY APPLN. INFO.: GB 2002-29281 Α 20021216 EP 2003-789283 A3 20031215 WO 2003-EP14263 W 20031215

OTHER SOURCE(S):

MARPAT 141:71530

GΙ

AB The title compds. [I; Rl = aryl having up to 10 carbon atoms; NR2R3 = heterocyclyl having up to 10 ring atoms and having 1-4 heteroatoms in the ring system; in free or salt form] which are useful for treating conditions mediated by of phosphodiesterase type 4 or the down-regulation or inhibition of TNF-α release, particularly obstructive or inflammatory airways diseases, were prepared E.g., a 3-step synthesis of 3-[6-(3-hydroxypyrrolidin-1-yl)-[1,7]naphthyridin-8-yl]benzonitrile, starting from 6-amino-8-bromo-1,7-naphthyridine and 3-cyanophenylboronic acid, which showed IC50 of 1 nM for inhibition of PDE4D, was given. Pharmaceutical compns. that contain compds. I and processes for preparing the compds. I are claimed.

IT 713145-28-9P 713145-30-3P 713145-47-2P

Ι

713145-56-3P 713145-62-1P

RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(preparation of [1,7]naphthyridines as PDE4 inhibitors)

RN 713145-28-9 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3-fluorophenyl)-1,7-naphthyridin-6-yl]-, ethyl ester (CA INDEX NAME)

RN 713145-30-3 CAPLUS

RN 713145-47-2 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-[3-(methylthio)phenyl]-1,7-naphthyridin-6-yl]- (CA INDEX NAME)

MeS N

RN 713145-56-3 CAPLUS

CN Acetic acid, [[1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-4-piperidinyl]oxy]- (9CI) (CA INDEX NAME)

RN 713145-62-1 CAPLUS

CN 4-Piperidineacetic acid, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]- (CF INDEX NAME)

IT 713145-07-4P 713145-08-5P 713145-09-6P 713145-10-9P 713145-11-0P 713145-12-1P 713145-13-2P 713145-14-3P 713145-15-4P 713145-16-5P 713145-17-6P 713145-18-7P 713145-19-8P 713145-20-1P 713145-21-2P 713145-22-3P 713145-23-4P 713145-24-5P 713145-25-6P 713145-26-7P 713145-27-8P 713145-29-0P 713145-31-4P 713145-32-5P 713145-33-6P 713145-34-7P 713145-35-8P 713145-36-9P 713145-37-0P 713145-38-1P 713145-39-2P 713145-40-5P 713145-41-6P 713145-42-7P 713145-43-8P 713145-44-9P 713145-45-0P 713145-46-1P 713145-48-3P 713145-49-4P 713145-50-7P 713145-63-2P 713145-64-3P 713145-65-4P 713145-66-5P 713145-67-6P 713145-68-7P 713145-69-8P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of [1,7] naphthyridines as PDE4 inhibitors) RN 713145-07-4 CAPLUS CN Benzonitrile, 3-[6-(3-hydroxy-1-pyrrolidinyl)-1,7-naphthyridin-8-yl]- (CA INDEX NAME)

RN 713145-08-5 CAPLUS
CN 1-Piperazinepropanenitrile, 4-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-,
monohydrochloride (9CI) (CA INDEX NAME)

● HCl

RN 713145-09-6 CAPLUS
CN 4-Piperidinecarboxylic acid, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-,
lithium salt (9CI) (CA INDEX NAME)

● Li

● HCl

RN 713145-11-0 CAPLUS CN 4-Piperidinecarboxamide, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]- (CA INDEX NAME)

RN 713145-12-1 CAPLUS

CN Benzonitrile, 3-[6-(4-hydroxy-1-piperidinyl)-1,7-naphthyridin-8-yl]- (CA INDEX NAME)

RN 713145-13-2 CAPLUS

CN Benzonitrile, 3-[6-[4-(hydroxymethyl)-1-piperidinyl]-1,7-naphthyridin-8-yl]- (CA INDEX NAME)

RN 713145-14-3 CAPLUS

CN Benzonitrile, 3-[6-[4-(2-hydroxyethyl)-1-piperidinyl]-1,7-naphthyridin-8-yl]- (CA INDEX NAME)

RN 713145-15-4 CAPLUS

CN Benzonitrile, 3-[6-[(2S)-2-(hydroxymethyl)-1-pyrrolidinyl]-1,7-naphthyridin-8-yl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 713145-16-5 CAPLUS

CN Piperazine, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-4-[(tetrahydro-2-furanyl)carbonyl]- (9CI) (CA INDEX NAME)

RN 713145-17-6 CAPLUS

CN 3-Piperidinecarboxamide, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]- (CA INDEX NAME)

RN 713145-18-7 CAPLUS

CN Benzonitrile, 3-[6-(4-morpholinyl)-1,7-naphthyridin-8-yl]- (CA INDEX NAME)

RN 713145-19-8 CAPLUS

CN 2-Pyrrolidinecarboxamide, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-, (2S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 713145-20-1 CAPLUS

CN L-Proline, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 713145-21-2 CAPLUS

CN Benzonitrile, 3-[6-(4-methyl-1-piperazinyl)-1,7-naphthyridin-8-yl]- (CA INDEX NAME)

RN 713145-22-3 CAPLUS

CN 1-Piperazineacetic acid, 4-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-, lithium salt (9CI) (CA INDEX NAME)

T.i

RN 713145-23-4 CAPLUS

CN 1-Piperazineacetic acid, 4-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-, ethyl ester (CA INDEX NAME)

RN 713145-24-5 CAPLUS

CN Piperazine, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-4-(methylsulfonyl)-(9CI) (CA INDEX NAME)

RN 713145-25-6 CAPLUS

CN Benzonitrile, 3-[6-(3,5-dimethyl-1-piperazinyl)-1,7-naphthyridin-8-yl]-, monohydrochloride (9CI) (CA INDEX NAME)

HCl

RN 713145-26-7 CAPLUS

CN Benzonitrile, 3-[6-(4-ethyl-1-piperazinyl)-1,7-naphthyridin-8-yl]- (CA INDEX NAME)

RN 713145-27-8 CAPLUS

CN 3-Azetidinecarboxylic acid, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-(CA INDEX NAME)

RN 713145-29-0 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3-fluorophenyl)-1,7-naphthyridin-6-yl]-, sodium salt (9CI) (CA INDEX NAME)

● Na

RN 713145-31-4 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(5-fluoro-2-methoxyphenyl)-1,7-naphthyridin-6-yl]-, potassium salt (9CI) (CA INDEX NAME)

K

RN 713145-32-5 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3-chlorophenyl)-1,7-naphthyridin-6-yl]-, ethyl ester (CA INDEX NAME)

RN 713145-33-6 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(5-chloro-2-methoxyphenyl)-1,7-naphthyridin-6-yl]-, ethyl ester (CA INDEX NAME)

RN 713145-34-7 CAPLUS

CN Benzoic acid, 3-[6-(4-morpholinyl)-1,7-naphthyridin-8-yl]- (CA INDEX NAME)

RN 713145-35-8 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3-chlorophenyl)-1,7-naphthyridin-6-yl]-, sodium salt (9CI) (CA INDEX NAME)

Na

RN 713145-36-9 CAPLUS

CN 3-Piperidinecarboxylic acid, 1-[8-(3-chlorophenyl)-1,7-naphthyridin-6-yl]-, ethyl ester (CA INDEX NAME)

RN 713145-37-0 CAPLUS

CN 3-Piperidinecarboxylic acid, 1-[8-(5-chloro-2-methoxyphenyl)-1,7-naphthyridin-6-yl]-, ethyl ester (CA INDEX NAME)

RN 713145-38-1 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(5-chloro-2-methoxyphenyl)-1,7-naphthyridin-6-yl]-, sodium salt (9CI) (CA INDEX NAME)

Na

RN 713145-39-2 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3-methoxyphenyl)-1,7-naphthyridin-6-yl]- (CA INDEX NAME)

RN 713145-40-5 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3,5-difluorophenyl)-1,7-naphthyridin-6-yl]- (CA INDEX NAME)

RN713145-41-6 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3-methylphenyl)-1,7-naphthyridin-6-yl]-(CA INDEX NAME)

RN

713145-42-7 CAPLUS Acetic acid, [[1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-4-CN piperidinyl]oxy]-, potassium salt (9CI) (CA INDEX NAME)

RN 713145-43-8 CAPLUS

4-Piperidinecarboxylic acid, 1-[8-(1,3-benzodioxol-5-yl)-1,7-naphthyridin-CN 6-yl]- (CA INDEX NAME)

RN 713145-44-9 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-[3-(trifluoromethoxy)phenyl]-1,7-naphthyridin-6-yl]- (CA INDEX NAME)

RN 713145-45-0 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3-chloro-4-fluorophenyl)-1,7-naphthyridin-6-yl]- (CA INDEX NAME)

RN 713145-46-1 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-4-methyl-, potassium salt (9CI) (CA INDEX NAME)

● K

RN 713145-48-3 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-[3-(methylsulfinyl)phenyl]-1,7-naphthyridin-6-yl]- (CA INDEX NAME)

RN 713145-49-4 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-(8-phenyl-1,7-naphthyridin-6-yl)- (CA INDEX NAME)

RN 713145-50-7 CAPLUS

CN 4-Piperidineacetic acid, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-, potassium salt (9CI) (CA INDEX NAME)

K

RN 713145-63-2 CAPLUS

CN 1-Piperazinepropanenitrile, 4-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-(CA INDEX NAME)

RN 713145-64-3 CAPLUS

CN Benzonitrile, 3-[6-(1-piperazinyl)-1,7-naphthyridin-8-yl]- (CA INDEX NAME)

RN 713145-65-4 CAPLUS

CN 1-Piperazineacetic acid, 4-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]- (CA INDEX NAME)

RN 713145-66-5 CAPLUS

CN Benzonitrile, 3-[6-(3,5-dimethyl-1-piperazinyl)-1,7-naphthyridin-8-yl]-(CA INDEX NAME)

RN 713145-67-6 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3-chlorophenyl)-1,7-naphthyridin-6-yl]-(CA INDEX NAME)

RN 713145-68-7 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(5-chloro-2-methoxyphenyl)-1,7-naphthyridin-6-yl]- (CA INDEX NAME)

RN 713145-69-8 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-4-methyl- (CA INDEX NAME)

IT 713145-51-8P 713145-52-9P 713145-55-2P

713145-57-4P 713145-58-5P 713145-59-6P

713145-60-9P 713145-61-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(Reactant or reagent)
 (preparation of [1,7]naphthyridines as PDE4 inhibitors)

RN 713145-51-8 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-, ethyl ester (CA INDEX NAME)

RN 713145-52-9 CAPLUS

CN 1-Piperazinecarboxylic acid, 4-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-, 1,1-dimethylethyl ester (CA INDEX NAME)

RN 713145-55-2 CAPLUS

CN Acetic acid, [[1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-4-piperidinyl]oxy]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

RN 713145-57-4 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-4-methyl-, ethyl ester (CA INDEX NAME)

RN 713145-58-5 CAPLUS

CN 4-Piperidinecarboxylic acid, 1-[8-[3-(methylthio)phenyl]-1,7-naphthyridin-6-yl]-, ethyl ester (CA INDEX NAME)

RN 713145-59-6 CAPLUS

CN Methanesulfonic acid, trifluoro-, 1-[8-[3-(methylthio)phenyl]-1,7-naphthyridin-6-yl]-4-piperidinyl ester (9CI) (CA INDEX NAME)

RN 713145-60-9 CAPLUS

CN 4-Piperidinamine, 1-[8-[3-(methylthio)phenyl]-1,7-naphthyridin-6-yl]- (CA INDEX NAME)

RN 713145-61-0 CAPLUS

CN 4-Piperidineacetic acid, 1-[8-(3-cyanophenyl)-1,7-naphthyridin-6-yl]-, ethyl ester (CA INDEX NAME)

REFERENCE COUNT:

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L1 STRUCTURE UPLOADED

L2 1 S L1

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L4 1 S L3 FULL

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                 LMEDLINE coverage updated
                 SCISEARCH enhanced with complete author names
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         JUL 02
NEWS
         JUL 02
                 CHEMCATS accession numbers revised
         JUL 02 CA/CAplus enhanced with utility model patents from China
NEWS
NEWS 6
         JUL 16 Caplus enhanced with French and German abstracts
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         JUL 18
                 CA/CAplus patent coverage enhanced
                 USPATFULL/USPAT2 enhanced with IPC reclassification
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                 FSTA enhanced with new thesaurus edition
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                 patents
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         AUG 20
                 CA/CAplus enhanced with CAS indexing in pre-1907 records
NEWS 14
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                 Full-text patent databases enhanced with predefined
                  patent family display formats from INPADOCDB
NEWS 15
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                 USPATOLD now available on STN
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                 CAS REGISTRY enhanced with additional experimental
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                 STN AnaVist, Version 2.0, now available with Derwent
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         SEP 13
                 INPADOCDB enhanced with monthly SDI frequency
         SEP 17
NEWS 20
                  CA/CAplus enhanced with printed CA page images from
                  1967-1998
NEWS 21
         SEP 17
                 CAplus coverage extended to include traditional medicine
                 patents
NEWS 22 SEP 24
                  EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS 23
         OCT 02
                 CA/CAplus enhanced with pre-1907 records from Chemisches
                  Zentralblatt
NEWS 24
         OCT 19
                 BEILSTEIN updated with new compounds
NEWS 25
         NOV 15
                 Derwent Indian patent publication number format enhanced
NEWS 26 NOV 19
                 WPIX enhanced with XML display format
              19 SEPTEMBER 2007: CURRENT WINDOWS VERSION IS V8.2,
NEWS EXPRESS
              CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007.
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FULL ESTIMATED COST

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=> s phosphodiesterase

27518 PHOSPHODIESTERASE

2939 PHOSPHODIESTERASES

L1 28110 PHOSPHODIESTERASE

(PHOSPHODIESTERASE OR PHOSPHODIESTERASES)

=> s l1 and inflammatory

191641 INFLAMMATORY

348 INFLAMMATORIES

191749 INFLAMMATORY

(INFLAMMATORY OR INFLAMMATORIES)

L2 1762 L1 AND INFLAMMATORY

=> s 12 and inhibit?

1980762 INHIBIT?

L3 1674 L2 AND INHIBIT?

=> s 13 and isoenzym?

66308 ISOENZYM?

L4 151 L3 AND ISOENZYM?

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578265 ASSAY?

L5 15 L4 AND ASSAY?

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L5 ANSWER 1 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2007:789095 CAPLUS

DOCUMENT NUMBER: 147:181512

TITLE: Screening for regulators of intracellular calcium

levels for control of NFAT transcription factors

INVENTOR(S): Rao, Anjana; Feske, Stefan; Hogan, Patrick; Gwack,

Yousang

PATENT ASSIGNEE(S): Cbr Institute for Biomedical Research, Inc., USA

SOURCE: PCT Int. Appl., 138pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PAT	ENT :	NO.			KIND DATE			i	APPL:	ICAT:		DATE						
		 2007 2007				A2 20070719 A9 20070907			. 1	WO 2	007-1		20070105						
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		RW:	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	ΓI,	FR,	GB,	GR,	HU,	IE,	
			IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,	
			CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	GH,	
			GM,	ΚE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,	
			KG,	ΚZ,	MD,	RU,	ТJ,	TM,	ΑP,	EA,	EP,	OA							
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	calcineurin inhibitors are described. The drugs target the											the							
	system of calcium uptake that regu													ds. that affect					
	intracellular calcium levels can be										ayed	by 1	thei	r ef	fect	s on			
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calcineurin inhibitors are described. The drugs target the system of calcium uptake that regulates calcineurin. Compds. that affect intracellular calcium levels can be assayed by their effects on NFAT, e.g. by use of an NFAT-dependent reporter gene, or by measuring NFAT binding to its binding site. Methods of measuring NFAT levels can also be used to diagnose disease including unexplained immunodeficiency. Alternatively, other calcium entry-mediated processes can be used as markers in screening. The role of calcium transporters is identified by a combination of mapping of genes associated with severe combined immunodeficiency in humans, and RNAi screening for effectors of calcium levels and NFAT nuclear transport in Drosophila. Human homologs of these genes were then identified.

L5 ANSWER 2 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2007:333039 CAPLUS

DOCUMENT NUMBER: 146:358872

TITLE: Pyrano[2,3-d]pyrimidines as nicotinic acid receptor

agonists for the treatment of dyslipidemia and their

preparation and pharmaceutical compositions

INVENTOR(S): Palani, Anandan; Su, Jing; Xiao, Dong; Huang, Xianhai;

Rao, Ashwin U.; Chen, Xiao; Tang, Haiqun; Qin, Jun; Huang, Ying R.; Aslanian, Robert G.; McKittrick, Brian A.; Degrado, Sylvia J.

PATENT ASSIGNEE(S):

USA

SOURCE: U

U.S. Pat. Appl. Publ., 239pp., Cont.-in-part of U.S.

Ser. No. 432,133.

CODEN: USXXCO

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE		
				-			
US 2007066630	A1	20070322	US 2006-600216		20061115		
US 2006264489	A1	20061123	US 2006-432133		20060511		
PRIORITY APPLN. INFO.:			US 2005-681848P	P	20050517		
			US 2005-715565P	P	20050909		
			US 2005-731039P	P	20051028		
			US 2006-432133	A2	20060511		

OTHER SOURCE(S):

MARPAT 146:358872

GI

AB A compound having the general structure of formula I: Formula I or a pharmaceutically acceptable salt, solvate, ester, or tautomer thereof, or a pharmaceutically acceptable salt, solvate, ester, or tautomer thereof, are useful in treating diseases, disorders, or conditions such as metabolic syndrome and dyslipidemia. Compds. of formula I wherein dotted line represents a single or double bond; if dotted line between X and CR7R7' is single, X is O and NH and derivs.; if dotted line between X and CR7R7' is double X is N; R1 is H, (un) substituted alkyl, alkenyl, alkynyl, haloalkyl, cycloalkyl, etc.; R2 is H, halo, (un) substituted alkyl, haloalkyl, etc.; R1R2 together with the atoms they are attached to may form a 5- to 6-membered (hetero)cycloalkenyl ring; R3 is absent, H, (un) substituted alkyl, cycloalkyl, (hetero) aryl, etc.; R4' and R6' are absent when dotted lines to the nitrogens are double bond and R4 and R6 are independently H, alkyl, halo, OH and derivs., NH2 and derivs., etc.; if dotted lines are single bond, R4R4' and R6R6' taken together is =0; R5 is absent, H, alkyl, alkynyl, alkylene-CO2-alkyl, etc.; R7R7' taken together is =0, when dotted line to X is single bond; R7 and R7' is H, alkyl, and (hetero)aryl; R7' is absent when dotted line to X is double bond; R7 is OH and derivs.; and their pharmaceutically acceptable salts, solvates, esters, and tautomers are claimed. Example compound II was prepared by cyclization of Me propionylacetate with barbituric acid. All the invention compds. were evaluated for their nicotinic acid receptor agonistic activity. From the assay, it was determined that several compds. exhibited cAMP EC50 value of 100 nM or less.

L5 ANSWER 3 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2006:1225392 CAPLUS

DOCUMENT NUMBER:

146:7973

TITLE:

Pyrano[2,3-d]pyrimidines as nicotinic acid receptor agonists for the treatment of dyslipidemia and their

preparation and pharmaceutical compositions

Palani, Anandan; Su, Jing; Xiao, Dong; Huang, Xianhai; Rao, Ashwin U.; Chen, Xiao; Tang, Haiqun; Qin, Jun;

Huang, Ying; Aslanian, Robert G.; Mckittrick, Brian

PATENT ASSIGNEE(S): Schering Corporation, USA SOURCE: PCT Int. Appl., 213pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

INVENTOR(S):

PAT	CENT	NO.			KIND DATE			i	APPL	ICAT:		DATE					
	2006 2006				A2 A3	A2 20061123 A3 20070308			1	WO 2	006-1		20060511				
	W: AE, AG, AL, CN, CO, CR, GE, GH, GM, KZ, LC, LK, MZ, NA, NG, SG, SK, SL, VN, YU, ZA,			CU, HR, LR, NI, SM,	CZ, HU, LS, NO, SY,	DE, ID, LT, NZ,	DK, IL, LU, OM,	DM, IN, LV, PG,	DZ, IS, LY, PH,	EC, JP, MA, PL,	EE, KE, MD, PT,	EG, KG, MG, RO,	ES, KM, MK, RU,	FI, KN, MN, SC,	GB, KP, MW, SD,	GD, KR, MX, SE,	
PRIORITY	VN, YU, ZA, RW: AT, BE, BG, IS, IT, LT, CF, CG, CI, GM, KE, LS, KG, KZ, MD, TY APPLN. INFO.:				CH, LU, CM, MW,	CY, LV, GA, MZ,	MC, GN, NA,	NL, GQ,	PL, GW, SL,	PT, ML,	RO, MR, TZ, 005-	SE, NE, UG, 6818	SI, SN, ZM, 48P 65P	SK, TD, ZW,	TR, TG, AM,	BF, BW,	BJ, GH, BY, 517

OTHER SOURCE(S):

MARPAT 146:7973

GΙ

A compound having the general structure of formula I: Formula I or a pharmaceutically acceptable salt, solvate, ester, or tautomer thereof, or a pharmaceutically acceptable salt, solvate, ester, or tautomer thereof, are useful in treating diseases, disorders, or conditions such as metabolic syndrome and dyslipidemia. Compds. of formula I wherein dotted line represents a single or double bond; if dotted line between X and CR7R7' is single, X is O and NH and derivs.; if dotted line between X and CR7R7' is double X is N; R1 is H, (un) substituted alkyl, alkenyl, alkynyl, haloalkyl, cycloalkyl, etc.; R2 is H, halo, (un) substituted alkyl, haloalkyl, etc.; R1R2 together with the atoms they are attached to may form a 5- to 6-membered (hetero)cycloalkenyl ring; R3 is absent, H, (un) substituted alkyl, cycloalkyl, (hetero) aryl, etc.; R4' and R6' are absent when dotted lines to the nitrogens are double bond and R4 and R6 are independently H, alkyl, halo, OH and derivs., NH2 and derivs., etc.; if dotted lines are single bond, R4R4' and R6R6' taken together is =0; R5 is absent, H, alkyl, alkynyl, alkylene-CO2-alkyl, etc.; R7R7' taken together is =0, when dotted line to X is single bond; R7 and R7' is H,

alkyl, and (hetero)aryl; R7' is absent when dotted line to X is double bond; R7 is OH and derivs.; and their pharmaceutically acceptable salts, solvates, esters, and tautomers are claimed. Example compound II was prepared by cyclization of Me propionylacetate with barbituric acid. All the invention compds. were evaluated for their nicotinic acid receptor agonistic activity. From the assay, it was determined that several compds. exhibited cAMP EC50 value of 100 nM or less.

ANSWER 4 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2006:607322 CAPLUS

DOCUMENT NUMBER: 145:241274

TITLE: The effects of a novel phosphodiesterase 7A

and -4 dual inhibitor, YM-393059, on

T-cell-related cytokine production in vitro and in

vivo

AUTHOR(S): Yamamoto, Satoshi; Sugahara, Shingo; Naito, Ryo;

Ichikawa, Atsushi; Ikeda, Ken; Yamada, Toshimitsu;

Shimizu, Yasuaki

CORPORATE SOURCE: Pharmacology Research Laboratories, Astellas Pharma

Inc., Ibaraki, 305-8585, Japan

SOURCE: European Journal of Pharmacology (2006), 541(1-2),

106-114

CODEN: EJPHAZ; ISSN: 0014-2999

PUBLISHER: Elsevier B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

YM-393059, (±)-N-(4,6-dimethylpyrimidin-2-yl)-4-[2-(4-methoxy-3methylphenyl)-5-(4-methylpiperazin-1-yl)-4,5,6,7-tetrahydro-1H-indol-1-

yl]benzenesulfonamide difumarate, is a novel phosphodiesterase

(PDE) inhibitor that inhibited the PDE7A

isoenzyme with a high potency (IC50 = 14 nM) and PDE4 with a moderate potency (IC50 = 630 nM). In a cell-based assay, YM-393059 was found to inhibit anti-CD3 antibody, Staphylococcal enterotoxin B, and phytohaemagglutinin-induced interleukin (IL)-2 production

in mouse splenocytes with IC50 values ranging from 0.48 to 1.1 μM . It also inhibited anti-CD3 antibody-induced interferon

(IFN)- γ and IL-4 production in splenocytes with IC50 values of 1.8 and 2.8 µM, resp. YM-393059's inhibition of anti-CD3

antibody-stimulated cytokine (IL-2, IFN-\gamma, and IL-4) production was 20to 31-fold weaker than that of YM976, a selective PDE4 inhibitor

However, orally administered YM-393059 and YM976 inhibited anti-CD3 antibody-induced IL-2 production equipotently in mice. In addition, YM-393059 inhibited lipopolysaccharide-induced tumor necrosis factor- α production in vivo more potently than IL-2 (ED50 values of 2.1

mg/kg and 74 mg/kg). In contrast to YM976, YM-393059 did not shorten the duration of α 2-adrenoceptor agonist-induced sleep in mice, which is a model for the assessment of the typical side effects caused by PDE4 inhibitors, nausea and emesis. YM-393059 is a novel and

attractive compound for the treatment of a wide variety of T-cell-mediated diseases.

48

ANSWER 5 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2005:447673 CAPLUS

DOCUMENT NUMBER: 143:20875

REFERENCE COUNT:

TITLE: Differentially expressed gene profile for diagnosing

and treating mental disorders

INVENTOR(S):

Akil, Huda; Atz, Mary; Bunney, William E., Jr.; Choudary, Prabhakara V.; Evans, Simon J.; Jones, Edward G.; Li, Jun; Lopez, Juan F.; Myers, Richard; Thompson, Robert C.; Tomita, Hiroaki; Vawter, Marquis

THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

P.; Watson, Stanley

PATENT ASSIGNEE(S): The Board of Trustees of the Leland Stanford Junior

University, USA

SOURCE: PCT Int. Appl., 226 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

1 ·

PATENT INFORMATION:

PATENT NO. DATE APPLICATION NO. KIND DATE _____ -----____ -----_____ A2 20050526 WO 2004-US36784 WO 2005046434 20041105 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG US 2005209181 A1 20050922 US 2004-982556 20041104 AU 2004-289247 AU 2004289247 **A**1 20050526 20041105 CA 2543811 **A**1 20050526 CA 2004-2543811 20041105 EP 1680009 A2 20060719 EP 2004-800741 20041105 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR, IS, YU PRIORITY APPLN. INFO.: US 2003-517751P P 20031105 A 20041104 US 2004-982556 W 20041105

The present invention provides methods for diagnosing mental disorders AB (e.g., psychotic disorders such as schizophrenia). The present invention uses DNA microarray anal. to demonstrate differential expression of genes in selected regions of post-mortem brains from patients diagnosed with mental disorders in comparison with normal control subjects. The invention also provides methods of identifying modulators of such mental disorders as well as methods of using these modulators to treat patients suffering from such mental disorders.

ANSWER 6 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2005:78243 CAPLUS

DOCUMENT NUMBER:

142:155827

TITLE:

Preparation of N-(cis-4-aminocyclohexyl)-2-(benzothienyloxy) nicotinamide derivatives as

inhibitors of 3',5'-cyclic nucleotide

phosphodiesterase 4 (PDE4)

INVENTOR(S):

Smith, Mya Coral Helen; Watson, Christine Anne Louise

WO 2004-US36784

Pfizer Inc, UK PATENT ASSIGNEE(S):

SOURCE:

U.S. Pat. Appl. Publ., 23 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2005020639	A1	20050127	US 2004-896112	20040720
US 7132435 CA 2536383	B2 A1	20061107 20050203	CA 2004-2536383	20040713

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WO 2005009438
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                                                                    20060125
    US .2007066645
                          Α1
                                20070322
                                            US 2006-555931
                                                                    20061102
PRIORITY APPLN. INFO.:
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OTHER SOURCE(S):
                         CASREACT 142:155827; MARPAT 142:155827
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AB This invention relates to nicotinamide derivs. of general formula (I) [R1 = H, halo, C1-4 alkyl; X = CH2, Y = S; or X = S and Y = CH2; Z = CO, SO2; R2 = each (un)substituted Ph, benzyl, naphthyl, heteroaryl or C3-8 cycloalkyl] or pharmaceutically acceptable salts or solvates thereof. These compds. are inhibitors of 3',5'-cyclic nucleotide phosphodiesterases (PDEs), i.e., PDE4A, PDE4B, PDE4C, and PDE4D which are isoforms or subtypes of the PDE4 isoenzyme family. They are particularly useful for the treatment of a great number of inflammatory, respiratory, and allergic diseases, disorders or conditions and for wounds and some of them are in clin. development mainly for treatment of asthma, chronic obstructive lung disease (COPD), bronchitis, and emphysema. Thus, cis-N-(4-aminocyclohexyl)-2-(2,3dihydrobenzo[b]thiophen-6-yloxy)-5-fluoronicotinamide (150 mg, 0.39 mmol), imidazo[1,2-a]pyridine-8-carboxylic acid (87 mg, 0.43 mmol), 1-hydroxybenzotriazole hydrate (58 mg, 0.43 mmol), 1-(3dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (82 mg, 0.43 mmol) and 4-methylmorpholine (47 μ L, 0.43 mmol) were dissolved in CH2Cl2 (20 mL) and the reaction mixture was stirred at room temperature for 18 h and was concentrated in vacuo. The residue was dissolved in DMF (10 mL) and stirred at room temperature for 18 h to give, after workup and silica gel chromatog., 130 mg (63%) imidazo[1,2-a]pyridine-8-carboxylic acid [cis-4-[[[2-[(2,3-a]pyridine-8-carboxylic acid [cis-4-[[[2-[(2,3-[a]pyridine-8-carboxylic acid [cis-4-[[[2-[(2,3-[a]pyridine-8-carboxylic acid [cis-4-[[[2-[(2,3-[a]pyridine-8-carboxylic acid [cis-4-[[[2-[(2,3-[a]pyridine-8-carboxylic acid [cis-4-[[[2-[(2,3-[a]pyridine-8-carboxylic acid [cis-4-[[a]pyridine-8-[a]pyrididihydrobenzo[b]thiophen-6-yl)oxy]-5-fluoropyridin-3-yl]carbonyl]amino]cyclohexyl]amide (II). Antiinflammatory properties of the nicotinamide

derivs. I were demonstrated by their ability to inhibit $TNF\alpha$ release from human peripheral blood mononuclear cells. II

showed IC50 of 0.6 nM in the above assay.

REFERENCE COUNT: 18 THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 7 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:60760 CAPLUS

Correction of: 2004:1036573

DOCUMENT NUMBER: 142:153477

Correction of: 142:16776

TITLE: Gene expression profiles and biomarkers for the

detection of Chagas disease and other disease-related

gene transcripts in blood

INVENTOR(S):
Liew, Choong-Chin

PATENT ASSIGNEE(S): Chondrogene Limited, Can.

33

SOURCE: U.S. Pat. Appl. Publ., 154 pp., Cont.-in-part of U.S.

Ser. No. 802,875.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

L5

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
US 2004241729	A1	20041202	US 2004-813097		20040330
US 2004014059	A1	20040122	US 2002-268730		20021009
US 2007031841	A1	20070208	US 2003-601518		20030620
US 2006134635	A1	20060622	US 2004-802875		20040312
US 2005191637	A1	20050901	US 2004-803737		20040318
US 2005196762	A1	20050908	US 2004-803759		20040318
US 2005196763	A 1	20050908	US 2004-803857		20040318
US 2005196764	A 1	20050908	US 2004-803858		20040318
US 2005208505	A 1	20050922	US 2004-803648		20040318
PRIORITY APPLN. INFO.:		•	US 1999-115125P	P	19990106
			US 2000-477148	B1	20000104
			US 2002-268730	A2	20021009
			US 2003-601518	A2	20030620
			US 2004-802875	A2	20040312
			US 2001-271955P	P	20010228
			US 2001-275017P	P	20010312
			US 2001-305340P	P	20010713
			US 2002-85783	A2	20020228
AR The present inventi	on ic	directed to	detection and measur	romoni	- of ~opo

The present invention is directed to detection and measurement of gene AB transcripts and their equivalent nucleic acid products in blood. Specifically provided is anal. performed on a drop of blood for detecting, diagnosing, and monitoring diseases, and in particular Chaqas disease, using gene-specific and/or tissue-specific primers. Affymetrix Human Genome U133 and ChondroChip microarrays were used to detect differentially expressed gene transcripts in hypertension, obesity, allergy, systemic steroids, coronary artery disease, diabetes type 2, hyperlipidemia, lung disease, bladder cancer, rheumatoid arthritis, osteoarthritis, liver cancer, schizophrenia, Chagas disease, asthma, and manic depression syndrome. The present invention describes methods by which delineation of the sequence and/or quantitation of the expression levels of disease-specific genes allows for an immediate and accurate diagnostic/prognostic test for disease or to assess the effect of a particular treatment regimen. [This abstract record is one of 3 records for this document necessitated by the large number of index entries required to fully index the document and publication system constraints.].

ACCESSION NUMBER: 2004:413097 CAPLUS

DOCUMENT NUMBER: 140:402343

TITLE: Diagnostics, drug screening and therapeutics for

diseases associated with human phosphodiesterase 4A (PDE4A)

INVENTOR(S): Golz, Stefan; Brueggemeier, Ulf; Geerts, Andreas

PATENT ASSIGNEE(S): Bayer Healthcare A.-G., Germany

SOURCE: PCT Int. Appl., 132 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	rent :	NO.			KIND DATE			j	APPL:	ICAT:		DATE						
WO	2004	0420	76		A2	_	2004	0521	1	WO 2	003-	EP11	879		2	0031		
WO	2004	0420	76		A3		2004	1014										
	W:	ΑE,	AG,	AL,	AM,	AT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
		co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	GE,	
		HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	ΚZ,	LC,	LK,			
	LR, LS, LT						MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,	NZ,	
	OM, PG, PH					PT,	RO,	RU,	SC,	SD,	SĖ,	SG,	SK,	SL,	SY,	TJ,	TM,	
		TN,	TR,	TT,	TZ,	UA,	UG,	US,	UŻ,	VC,	VN,	YU,	ZA,	ZM,	ZW			
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,	
		KG,	ΚZ,	MD,	RU,	ТJ,	TM,	ΑT,	ΒĒ,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	
		FI,	FR,	GB,	GR,	HU,	ΙE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,	TR,	
		BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG	
AU	AU 2003274082									AU 2	2740							
PRIORITY	PRIORITY APPLN. INFO.:									EP 20	002-	2499	4	7	A 20	0021	108	
									Ţ	WO 21	003-1	EP11	879	7	W 2	0031	025	

AB The invention provides a human PDE4A which is associated with the disorders of the peripheral and central nervous system, cardiovascular diseases, hematol. diseases, inflammation, gastroenterol. diseases and endocrinol. diseases. The cDNA sequence and the encoded amino acid sequence of PDE4A are disclosed. The expression profile of PDE4A in various human tissues is shown. The invention also provides assays for the drug screening and identification of compds. useful in the treatment or prevention of disorders of the peripheral and central nervous system, cardiovascular diseases, hematol. diseases, inflammation, gastroenterol. diseases and endocrinol. diseases. The invention also features compds. which bind to and/or activate or inhibit the activity of PDE4A as well as pharmaceutical compns. comprising such compds.

L5 ANSWER 9 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2004:388025 CAPLUS

DOCUMENT NUMBER: 140:385801

TITLE: KF19514, a phosphodiesterase 4 and 1

inhibitor, inhibits

TNF- α -induced GM-CSF production by a human bronchial epithelial cell line via inhibition

of PDE4

AUTHOR(S): Sasaki, K.; Manabe, H.

CORPORATE SOURCE: Pharmaceutical Research Institute, Kyowa Hakko Kogyo

Co., Ltd., Shizuoka, 411-8731, Japan

SOURCE: Inflammation Research (2004), 53(1), 31-37

CODEN: INREFB; ISSN: 1023-3830

PUBLISHER: Birkhaeuser Verlag

DOCUMENT TYPE: Journal LANGUAGE: English

AB Bronchial epithelium plays an important role in the regulation of inflammatory reactions in the airways. We investigated the effect of KF19514, a phosphodiesterase (PDE) 4 and 1 inhibitor

, on granulocyte-macrophage colony-stimulating factor (GM-CSF) production by a human bronchial epithelial cell line, BEAS-2B. BEAS-2B cells were stimulated with the tumor necrosis factor- α (TNF- α) and various concns. of test agents for 48 h. Supernatants were assayed for GM-CSF by using an ELISA. In addition, intracellular cAMP levels were measured in the presence of various agents. KF19514 significantly inhibited the release of GM-CSF by BEAS-2B cells in a concentration-dependent manner. The other PDE4 inhibitors and cAMP-elevating agents also inhibited the GM-CSF production In the BEAS-2B cells, KF19514 and PDE4 inhibitors concentration-dependently increased intracellular cAMP levels. The inhibitory effect of KF19514 on the GM-CSF production was significantly reduced by a cAMP-dependent protein kinase A (PKA) inhibitor, H89. Other PDE isoenzyme inhibitors did not inhibit the GM-CSF production by BEAS-2B cells, and did not elevate the intracellular cAMP levels. These results indicate that KF19514 and PDE4 inhibitors reduce TNF- α -induced GM-CSF production of BEAS-2B cells via a cAMP-dependent pathway. PDE4 may be a possible target for the regulation of cytokine production in epithelial cells.

REFERENCE COUNT:

40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 10 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2004:130977 CAPLUS

DOCUMENT NUMBER: 140:281023

TITLE: Anti-inflammatory potential of the selective

phosphodiesterase 4 inhibitor

N-(3,5-dichloro-pyrid-4-yl)-[1-(4-fluorobenzyl)-5-

hydroxy-indole-3-yl]-glyoxylic acid amide (AWD

12-281), in human cell preparations

Draheim, Regina; Egerland, Ute; Rundfeldt, Chris AUTHOR(S): CORPORATE SOURCE:

Departments of Pharmacology and Molecular Biology,

Elbion AG, Radebeul, Germany

Journal of Pharmacology and Experimental Therapeutics SOURCE:

(2004), 308(2), 555-563

CODEN: JPETAB; ISSN: 0022-3565

PUBLISHER: American Society for Pharmacology and Experimental

Therapeutics

DOCUMENT TYPE: Journal LANGUAGE: English

AWD 12-281 is a potent (IC50 = 9.7 nM) and highly selective inhibitor of the phosphodiesterase 4 (PDE4) isoenzyme with low affinity to the high-affinity rolipram-binding The compound was optimized for topical treatment of asthma, chronic obstructive pulmonary disease (COPD), and allergic rhinitis. the present study was to assess the effect of AWD 12-281 in human inflammatory cells. Peripheral blood mononuclear cells (PBMCs), diluted whole blood, and human nasal polyp cells derived from surgically resected nasal polyps from patients with polyposis comprise sources of target tissue cells that can be used to predict anti-inflammatory effects in patients. AWD 12-281 was capable of suppressing the production of cytokines in stimulated PBMCs: interleukin-2 (IL-2, phytohemagglutinin stimulation), IL-5 (Con A stimulation), IL-5 and IL-4 (anti-CD3/anti-CD28 co-stimulation), and lipopolysaccharide-stimulated release of tumor necrosis factor α (TNF α). The corresponding values for half-maximum inhibition, EC50, for AWD 12-281 were within a narrow range (46-121 nM). Comparing the effect of AWD 12-281 with roflumilast, cilomilast (SB 207499), rolipram (RPR-73401), and 1-(3-nitrophenyl)-3-(4pyridylmethyl)pyrido[2,3-d]pyrimidin-2,4(1H,3H)-dione (RS-25344-000), it could be shown that the PDE4 inhibitory activity was closely correlated with inhibitory potential as measured by the above-described assays. AWD 12-281 was also shown to suppress $TNF\alpha$ release in dispersed nasal polyps (EC50 = 111 nM) and in diluted

whole blood (EC50 = 934 nM). The reduced activity in human blood may be related to high plasma protein binding. Currently, phase II clin. studies are under way to evaluate the therapeutic potential of AWD 12-281 in asthma, COPD, and allergic rhinitis.

REFERENCE COUNT:

40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 11 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2002:601116 CAPLUS

DOCUMENT NUMBER:

137:351413

TITLE:

Potential role of phosphodiesterase 7 in

human T cell function: comparative effects of two

phosphodiesterase inhibitors

AUTHOR(S):

Nakata, A.; Ogawa, K.; Sasaki, T.; Koyama, N.; Wada, K.; Kotera, J.; Kikkawa, H.; Omori, K.; Kaminuma, O.

CORPORATE SOURCE:

Discovery Research Laboratory, Tanabe Seiyaku Co.

Ltd., Saitama, Japan

SOURCE:

Clinical and Experimental Immunology (2002), 128(3),

460-466

CODEN: CEXIAL; ISSN: 0009-9104 Blackwell Science Ltd.

PUBLISHER: Blackwe
DOCUMENT TYPE: Journal

DOCUMENT TYPE: LANGUAGE: Journal English

Even though the existence of phosphodiesterase (PDE) 7 in T cells has been proved, the lack of a selective PDE7 inhibitor has confounded an accurate assessment of PDE7 function in such cells. In order to elucidate the role of PDE7 in human T cell function, the effects of two PDE inhibitors on PDE7A activity, cytokine synthesis, proliferation and CD25 expression of human peripheral blood mononuclear cells (PBMC) were determined Recombinant human PDE7A was obtained and subjected to cAMP-hydrolysis assay. PBMC of Dermatophagoides farinae mite extract (Df)-sensitive donors were stimulated with the relevant antigen or an anti-CD3 monoclonal antibody (MoAb). PBMC produced IL-5 and proliferated in response to stimulation with Df, while stimulation with anti-CD3 MoAb induced CD25 expression and mRNA synthesis of IL-2, IL-4 and IL-5 in peripheral T cells. A PDE inhibitor, T-2585, which suppressed PDE4 isoenzyme with high potency (IC50 = 0.00013 μ m) and PDE7A with low potency (IC50 = 1.7 μ m) inhibited cytokine synthesis, proliferation and CD25 expression in the dose range at which the drug suppressed PDE7A activity. A potent selective inhibitor of PDE4 (IC50 = 0.00031 μ M), RP 73401, which did not effectively suppress PDE7A (IC50 > 10 μM), inhibited the Dfand anti-CD3 MoAb-stimulated responses only weakly, even at 10 $\mu\text{M}.$ PDE7 may play a critical role in the regulation of human T cell function, and thereby selective PDE7 inhibitors have the potential to be used to treat immunol. and inflammatory disorders.

REFERENCE COUNT:

50 THERE ARE 50 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 12 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2002:594822 CAPLUS

DOCUMENT NUMBER:

137:154857

TITLE:

Preparation of nicotinamide biaryl derivatives as

inhibitors of PDE4 isozymes

INVENTOR(S):

Chambers, Robert James; Magee, Thomas Victor; Marfat,

Anthony

PATENT ASSIGNEE(S): SOURCE:

Pfizer Products Inc., USA PCT Int. Appl., 224 pp.

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA?	CENT 1	NO.			KIND DATE				APPLICATION NO.							DATE		
WO	2002	0608	 75		A1	_	2002	8080		WO	2001-	IB23	 41			20013	206	
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BE	3, BG,	BR,	BY,	ΒZ,	CA	, СН,	CN,	
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC	, EE,	ES,	FI,	GB,	GD	, GE,	GH,	
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		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	M	, MW,	MX,	MZ,	NO,	ΝZ	, OM,	PH,	
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SF	K, SL,	ТJ,	TM,	TR,	TT	, TZ,	UA,	
		UG,	US,	UZ,	VN,	YU,	ZA,	ZW										
	RW:										, TZ,							
		KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	CF	I, CY,	DE,	DK,	ES,	FI	, FR,	GB,	
		GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	TF	R, BF,	ВJ,	CF,	CG,	CI	, CM,	GΑ,	
		GN,	GQ,				NE,	SN,	TD,	TO	;							
	2436				A1						2001-							
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EP	1355										2001-					20011		
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							RO,	•			•							
	2003		0		Α						2003-					20011		
BR	2001	0168	52		Α		2004	0225		BR	2001-	1685	2			20011		
HU	2004	0006	37		A2		2004	0628		HU	2004-	637				20011		
JP	2004 1518	5203	86		\mathbf{T}		2004			JΡ	2002-	5610	26			20011		
CN	1518	542			Α		2004			CN	2001-	8230	71			20011		
NZ	5264	53			Α		2005			ΝZ	2001-	5264	53			20011		
	2002		12		AI		2002			US	2002-	6281	3			20020	131	
	6649				B2		2003						_					
	2003						2005				2003-							
	2003				A		2004			ZA	2003-	4894				20030	1624	
	2004		03		A1		2004			US	2003-	6139	88			20030	702	
	6953				B2 A A		2005										.=	
	1080		0.7		A		2004			BG	2003- 2003-	1080	38			20030	1728	
NO	2003	0033	9/		A		2003			NO	2003-	3397	0.7					
	MX 2003FA00007				А		2003	1113		MX	2003	PAGE	8 / 02 D		_	20030	1/30	
PKIORIT	RIORITY APPLN. INFO.:									US	2001-	Z054	92P		۲ ت	20010	1737	
										WO	2002 2001	4201	4 ⊥ ວ		W	2002(.∠Ub	
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OTHER S	JUKCE	(5):			MARPAT 137:15485			⊃ <i>1</i>										

OTHER SOURCE(S): MARPAT 137:1

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB The title compds. [I; g = 0-1; j = 0-1; provided that when j = 0, n must be 2; k = 0-1; m = 0-2; n = 1-2; W1 = 0, Sot (t = 0-2), NR3; W2 = OCR9R10, or absent; Y = CR1, NOk (k = 0-1); R9, R10 = H, F, CF3, etc.; or R9 and R10 are taken together, but only in the case where m = 1, to form a spiro moiety; R7, R8 have the same meaning as R9, R10 except that one of them must be H; R1, R2 = H, F, Cl, etc.; R3 = H, alkyl, Ph, etc.; R4-R6 = H, F, Cl, etc.; Q1 = Ph, benzodioxyl, etc.; Q2 = biaryl moiety], useful as inhibitors of PDE4 in the treatment of diseases regulated by the activation and degranulation of eosinophils, especially asthma, chronic bronchitis, and chronic obstructive pulmonary disease, were prepared E.g., a multi-step synthesis of the amide II, starting from Me 3-bromobenzoate and 4-formylbenzeneboronic acid, was given. Compds. I showed anti-inflammatory activity at 0.0001 μM to 20.0 μM in whole blood assay for LTE4.

REFERENCE COUNT:

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ACCESSION NUMBER: 2001:136945 CAPLUS

DOCUMENT NUMBER: 134:193441

TITLE: Preparation of hypoxanthines and thiohypoxanthines as

phosphodiesterase IV inhibitors

INVENTOR(S): Chasin, Mark; Hofer, Peter; Cavalla, David

PATENT ASSIGNEE(S): Euro-Celtique S.A., Luxembourg

SOURCE: PCT Int. Appl., 68 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	rent :	NO.			KIND DATE				APPL	ICAT	ION 1	NO.	DATE				
WO	2001	0119	 67		A1		2001	0222		WO 2	000-	US21	 836		2	0000	809
	W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
							DM,										
		HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,
		LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,
		SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,
		YU,	ZA,	ŻW,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	TJ,	TM				
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	ŪG,	ZW,	AT,	BE,	CH,	CY,
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,
		CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	ΝĒ,	SN,	TD,	TG			
CA	2379	356			A1		2001	0222		CA 2	000-	2379	356		2	0000	809
	1202									EP 2	000-	9539	25		2	0000	809
EP	1202	628			В1		2004	1013									
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
							RO,										
JP	2003 2791	5064	67		T		2003	0218		JP 2	001-	5163	30		2	0000	809
AT	2791	13		•	T		2004	1015		AT 2	000-	9539	25		2	0000	809
PRIORIT	RIORITY APPLN. INFO.:									US 1999-148623P				P 19990812			
										WO 2000-US21836					W 20000809		
OTHER SO	OURCE	(S):			MAR:	PAT	134:	1934	41								

$$Q^{1} = (CH_{2})_{n}$$
 $Q^{2} = (CH_{2})_{n}$
 $Q^{2} = (CH_{2})_{n}$
 $Q^{2} = (CH_{2})_{n}$
 $Q^{2} = (CH_{2})_{n}$

AΒ Title compds. (I) [wherein R3 and R8 = independently (cyclo)alkyl, alkenyl, alkynyl, Q1, or Q2; R6 = S or O; n = 0-1; Z = a bond, CH2, NH, O, or S; A and B can form a ring by adding 1-3 CH2 groups when Z = CH2, NH, O or S; and A and B are not in a ring when Z = a bond, wherein A and B = . independently H, halo, (cyclo)alkyl, (cyclo)alkoxy, OH, or (un)substituted Ph, benzyl, or benzyloxy; L and M = independently H or Me; W = Q1, OH, (hetero)aryl, heterocyclyl, or (un)substituted benzyloxy] were prepared as selective phosphodiesterase (PDE) IV inhibitors. For example, amidation of 2-(4-fluorobenzyloxy)-2-methylpropionyl chloride with 5,6-diamino-1-(3,4-dimethoxybenzyl)-2-thiouracil using TEA in THF (20.4%), followed by cyclization with NaOH to form the 2-thioxanthine (79.1%) and treatment with Raney nickel in 1-propanol (67.2%), afforded the hypoxanthine (II). In assays measuring isolated PDE isoenzyme activity, II selectively inhibited PDE IV compared to PDE III and PDE V with IC50 values of 1.079 µM, 69.62 μM , and 35.80 μM , resp. As a result, I are expected to induce the desirable anti-asthmatic effects associated with PDE IV inhibition without causing the undesirable cardiovascular stimulation associated with PDE III inhibition (no data). I are useful in the treatment of asthma, allergies, inflammation, depression, dementia, and other disease states associated with abnormally high physiol. levels of cytokine (no data). REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS

L5 ANSWER 14 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1999:36647 CAPLUS

DOCUMENT NUMBER: 130:222068

TITLE: Phosphodiesterase 4B gene transcription is

activated by lipopolysaccharide and inhibited

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

by interleukin-10 in human monocytes

AUTHOR(S): Ma, Dongmin; Wu, Ping; Egan, Robert W.; Billah, M.

Motasim; Wang, Peng

CORPORATE SOURCE: Allergy Department, Schering-Plough Research

Institute, Kenilworth, NJ, USA

SOURCE: Molecular Pharmacology (1999), 55(1), 50-57

CODEN: MOPMA3; ISSN: 0026-895X

PUBLISHER: American Society for Pharmacology and Experimental

Therapeutics

DOCUMENT TYPE: Journal LANGUAGE: English

There are 4 different genes encoding the cAMP-specific phosphodiesterase (PDE4) isoenzymes (A, B, C, and D).

CAMP has been the only agent known to induce PDE4 gene expression. Here, the authors demonstrate, for the first time, that lipopolysaccharide (LPS) selectively stimulated PDE4B mRNA production in human monocytes. The LPS stimulation occurred very rapidly (in 30-45 min) and in a dose-dependent manner (0.01-100 ng/mL). The authors also demonstrate that LPS induction of PDE4B mRNA expression was inhibited strongly by interleukin (IL)-10. The inhibition with IL-10 was dose-dependent (0.1-10 ng/mL). IL-4 also inhibited the LPS induction, but to a lesser extent than IL-10. PDE4B mRNA expression was also stimulated by dibutyryl-cAMP. Interestingly, unlike LPS induction, the dibutyryl-cAMP induction of PDE4B mRNA expression was not inhibited by IL-10. By performing nuclear run-on and mRNA stability assays, the authors demonstrate further that IL-10 inhibited LPS-stimulated PDE4B mRNA synthesis by abolishing the gene transcription rather than by enhancing mRNA degradation Thus, PDE4B, as the only LPS-inducible PDE4 subtype, may be an appropriate target for discovering antiinflammatory drugs.

39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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1995:981130 CAPLUS ACCESSION NUMBER:

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TITLE: Effects of nonselective and isoenzyme

> selective cyclic nucleotide phosphodiesterase inhibitors on antigen-induced cytokine gene expression in peripheral blood mononuclear cells Essayan, David M.; Huang, Shau-Ku; Kagey-Sobotka,

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CORPORATE SOURCE: Division of Clinical Immunology, Johns Hopkins Asthma

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Cyclic nucleotide phosphodiesterase (PDE) enzymes may participate in regulation of the inflammatory response through their effects on second messengers. Here, the authors investigated the role of nonselective and isoenzyme selective PDE inhibitors in altering the antigen-driven cytokine gene expression of peripheral blood mononuclear cells (PBMCs) from atopic individuals. Ragweed and tetanus toxoid were used as model antigens. The nonselective PDE inhibitor, 3-isobutyl-1-methylxanthine (IBMX), and the selective PDE4 inhibitor, rolipram, markedly suppressed interleukin-5 (IL-5) and interferon γ (IFN γ) gene expression in both antigen-driven systems. Gene expression for IL-4 was unaffected by these agents in the ragweed-driven system. Message for IL-4 could not be detected in the tetanus toxoid-driven system, despite the use of a quant., competitive reverse transcription-polymerase chain reaction (RT-PCR) assay sensitive to <10 fg of target template. The PDE3 inhibitor, siguazodan, was ineffective in downregulating gene expression for the proinflammatory cytokines assay; when used in combination with the PDE4 inhibitor, the PDE3 inhibitor provided no increase in efficacy over that seen with the PDE4

inhibitor alone. Gene expression for the A and B isoforms of the PDE4 in PBMCs was unaffected by antigen stimulation or treatment with the PDE4 inhibitor; however, differences in expression of these 2 isoforms were apparent when a variety of immune cell lines were studied. These data support the hypothesis that the primary anti-inflammatory target for PDE inhibition in PBMCs is the PDE4. Furthermore, the expression of various isoforms of this enzyme may differ between immune cell types. Finally, PDE4 isoform expression in PBMCs is independent of treatment with an isoenzyme selective inhibitor.

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